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Origin of anthropogenic hydrocarbons and halocarbons measured in the summertime european outflow (on Crete in 2001)

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Abstract. During the Mediterranean Intensive Oxidant Study MINOS in August 2001, 87 air samples were collected at the ground-based station Finokalia (35°19'N, 25°40'E) on the north coast of Crete and subsequently analysed by GC-MS. The analysis includes various hydrocarbons, organo-halogens, HCFCs and CFCs. These compounds have a wide variety of sources and sinks and a large range of atmospheric lifetimes. We evaluated the characteristics of the sampling site in terms of proximity to individual sources by plotting the measured variability of these species against lifetime. The resulting linear relationship suggests that the sampling site is representative of intermediate conditions between a remote site and one that is in the vicinity of a wide variety of sources. Our analysis of air mass origin and chemical ratios also shows that several distinct anthropogenic sources influenced the atmospheric composition over Crete. Propane observations are compared to a global model to assess the fossil fuel related emission inventory. Although the model reproduces the general pattern of the propane variations, the model mixing ratios are systematically too low by a factor of 1.5 to 3, probably due to an underestimation of the propane emissions from east European countries in the underlying global database EDGAR. Another important finding was that methyl chloroform, a compound banned under the Montreal protocol, showed significant enhancements from background, which were well correlated with CFC-113. This suggests continued use and emission of methyl chloroform by one or more European countries. We also discuss the observed variations of methyl bromide and suggest that the significant peak observed on 12 August 2001 reflects heavy agricultural use as a soil fumigant in Italy.

1 Introduction

Volatile organic compounds (VOC) comprise a large number of chemical families, including alkanes, alkenes, carbonyls, alcohols, organo-halogens and CFCs. These compounds have a large range of lifetimes from a few hours for pentane to several hundred years for CFC-114.

The most reactive gases (non-methane hydrocarbons NMHCs- and partly oxygenated NMHCs) play an important role in the oxidising capacity of the atmosphere. Their atmospheric breakdown, in the presence of nitrogen oxides, leads to the photochemical build-up of ozone. Therefore, it is essential to determine accurately the budget and distribution of these compounds. While their sinks (mostly through reaction with OH, although the reactions with O₃ and NO₃ can also be significant for unsaturated NMHCs) are quite well quantified by laboratory studies, large uncertainties remain with respect to their sources. These compounds have a wide variety of biogenic sources (vegetation, ocean, etc) and anthropogenic sources (fossil fuel use and combustion, biomass burning, etc.), for which emissions still need to be confirmed, determined or quantified.

The fact that long-lived species (organo-halogens and CFCs) are radiatively active and that their potential to release chlorine or bromine in the stratosphere leads to ozone destruction, make them important species to study. Following the Montreal Protocol (1987) and subsequent amendments, the consumption of several halocarbons has now been phased out in developed countries. Several background studies have already reported the expected decrease of some of these compounds in the atmosphere and have also provided information about their regional sources. For instance, measurements of diverse halocarbons performed at Mace Head, Ireland, were used to estimate the intensity and distribution of their western European emissions (Ryall et al., 2001 and

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references therein). To our knowledge, no measurements have been performed directly in the outflow from eastern Europe in order to determine the eastern European contribution to their global budgets.

The MINOS campaign on Crete was dedicated to the observation of long-range transport of pollution and presented an ideal opportunity to perform measurements of VOCs in the outflow of eastern Europe and to study their origin. This paper focuses on observations of alkanes (propane, butanes, pentanes) and halocarbons (particularly methyl chloroform, CFC-113 and methyl bromide) and discusses their various anthropogenic emissions. A detailed analysis of the oxygenated compounds acetone and methanol is given elsewhere (Salisbury et al., 2003).

2 Experimental

2.1 Site and sampling

Air samples were collected at the atmospheric measurement station Finokalia 35°19'N, 25°40'E, 150 m above sea level, a remote site on the northern coast of Crete. Air samples were pressurised in 2.4 L electropolished stainless steel canisters previously flushed with synthetic air and then pressurised to about 4 bar. Canisters were made at the workshop of the Max Planck Institute and were equipped with stainless steel valves (Nupro SS4H). The canisters were not humidified but the quite high relative humidity observed during the MINOS campaign (63% on average) was considered to be sufficient to prevent losses associated with dry air. Using a Teflon membrane pump (KNF, Neuberger, Germany), canisters were flushed at least 5 times with ambient air and then pressurised to approximately 4 bar. Analysis of the samples was carried out in the laboratory in Mainz between 3 and 6 months after sampling. Stability of the measured compounds over this long period of time has been checked by filling two cylinders with the high-pressure working standard and by measuring them after 1, 3, and 5 months against the working standard. The working standard was also air collected from Finokalia (see Sect. 2.3). For all the compounds presented here but one, no drift in the concentrations was observed and the concentrations remained within the uncertainty range. The only exception was chloroform, which showed values lower than expected (up to 30%).

In August 2001, the site was influenced by on-shore north-north westerly winds with an average speed of 8 m s^{-1} . This wind advected pollutants from eastern Europe across the Aegean Sea to Crete (see Salisbury et al., 2003, for a map of the region and a complete description of air mass origin). Between 1 and 16 August 2001, 87 air samples were collected at six-hour intervals and at much shorter intervals on selected days.

2.2 Instrumentation

The analytical instrumentation consisted of a gas chromatograph-mass spectrometer system (GC-MS 6890/5973, Agilent Technology, Palo Alto, CA, USA) which has been modified from the commercial version for analysis of low-level ambient air samples. A sample preparation/cryoconcentrator unit was built (as detailed below) and the valves/heating system was automated using a custom built electronic programmable controller. To avoid water related problems (shifting of retention times or ice blocking the trap), water was removed by a magnesium perchlorate trap (replaced approximately every 100 measurements), commonly used for VOC analysis (Namiesnik and Wardencki, 1999 and references therein). Where possible, the lines and the switching valves were incorporated in a heated box ($\sim 50^\circ\text{C}$). The dryer, which was outside the box, was separately heated to above 75°C to improve drying efficiency and prevent condensation. A known volume of sample (about 1 liter) was cryogenically concentrated at -72°C using a Neslab cc-100 circulation cooler (Portsmouth, USA) at a flowrate of $30\text{--}40 \text{ ml min}^{-1}$ in a stainless steel microtrap (length: 30 cm, ID: 0.076 cm), packed with porous silica beads (Unibeads 1S, 80/100 mesh, Alltech) over a length of about 10 cm. The exact volume sampled was determined by the pressure difference in a known volume downstream of the trap, held at 50°C . For GC injection, the cold trap is heated to 200°C in less than 20 s. The carrier gas was Helium (99.9999, with a Supelco® catalytic purifier in-line) at an initial flow rate of 3.7 ml min^{-1} . A $60 \text{ m} \times 0.248 \text{ mm}$ DB-5 capillary column (J & W Scientific) was held at -65°C for one minute and then heated to 120°C at a rate of 8°C min^{-1} , followed by ramping to 175°C at a rate of $70^\circ\text{C min}^{-1}$ and remained at this temperature for 5 min before finally being heated at the same rate to 250°C and then maintained at this temperature for 6 min. A complete run lasts about 35 min and allows a good separation of more than 60 identified compounds. The mass spectrometer is operated in single ion mode (SIM) for maximum sensitivity of the measurements. More than half of the samples were measured twice and, on average, the two measurements agreed within 3.5% (standard deviation over mean).

2.3 Calibration and characteristics of the analysis system

Calibrations were performed against a working standard, a 10-L aluminium cylinder (Scott Marin, Riverside, California) filled at the Finokalia station two months before the campaign by using a three stage RIX oil-free piston compressor (Mak and Brenninkmeijer, 1994). This cylinder, along with a second, filled a couple of hours later, was then calibrated at the National Center for Atmospheric Research (NCAR, Boulder, Colorado) relative to a gravimetrically prepared standard referenced to standards by the National Institute of

Table 1. Average mixing ratios of selected VOCs measured during MINOS. The second column gives the corresponding monitored ions (the first ion was used for quantification). The clean air concentrations are an average of 4 samples collected on 7 August (see text). Mace Head data (August 2001 mean) are from the ALE/GAGE/ALGAGE network (<http://cdiac.ornl.gov/ndps/aleage.html>)

Compounds	Masses	Overall Air pptv	Clean Air pptv	Mace Head pptv
Propane	43, 41	235±121	123±21	
i-butane	43, 41	72±178	34.3±8	
n-butane	43	86±93	42.7±11	
i-pentane	43, 42, 41, 57	97±278	40±14	
n-pentane	43	36±64	17±4	
Benzene	78	101±56	50±14	
Methyl chloride	50, 52	588±104	537±38	
Dichloromethane	49, 84	32±23	25±3	
Chloroform	83, 47	14±3	11±1	10.49
Methyl-chloroform	97, 61	40±7	33.9±0.4	
CCl ₄	117, 119	93±7	92±2	95.76
Methyl bromide	94, 96	12±3	15±3	
CFC-12	85	530±18	507±8	545.42
CFC-113	101, 103, 151	82±7	76.2±0.8	
CFC-114	85, 87	13.4 0.5	12.8±0.2	
HCFC-22	51	158±25	140±9	
HCFC-142b	65	16±3	15.1±0.5	14.73

Values are reported as geometric means (to minimize skew errors of a non-normal distribution) with one standard deviation.

Standards and Technology (NIST) (estimated accuracy of 2–3%). Therefore, all the VOC measurements presented in this paper are calibrated on the NCAR calibration scale. Details of the NCAR calibration procedure are given in Schauffler et al. (2003) and references therein. About one year after the collection of these two cylinders, we measured our working standard against the other calibrated cylinder and we found the expected concentration (3.5% difference on average). Assuming the two cylinders (which were taken at different times of day) did not drift in exactly the same way, we may conclude that the concentrations in our working standard did not drift over the period of MINOS measurements. Two different calibration procedures were used. At first, the mass spectrometer was tuned every week and a calibration was performed several times at the beginning and at the end of a week. A calibration was once performed in the middle of the week and it was found that the decrease of MS sensitivity over the week was almost linear and therefore the response factors for every day were linearly interpolated over the week. However, as the decrease of the sensitivity was significant from week to week, it could not be extrapolated between weeks. Therefore, for the last 37 samples a second method was applied in which the working standard was analysed at the beginning and at the end of the day (the MS still being tuned each week). The average response factor of these two analyses was used to calibrate the samples measured on the same day. There was never a trend in the daily calibration greater than the mean precision which was 15% (see below).

At least two blanks (using the same analytical procedure but without collecting sample) were performed at the start of each measurement sequence and showed a generally clean baseline with only minor impurities of benzene, which were well within its measurement uncertainty.

Several tests were performed in the laboratory to evaluate the characteristics of the analytical system. Detection limits were in the range of 1–10 parts per trillion (10^{-12} mol per mol by volume, pptv) for the compounds reported here. Linearity of the analytical system was better than 3% for all the compounds presented. The linearity was determined by injecting different amounts (from 0.03 L to 1 L) of the same sample and by monitoring the corresponding peak area. The obtained linear fits were used to calculate the residuals. The residuals were calculated as the difference between the actual value and the linear fit (for each point between 0.03 and 1 L injections), these differences being then divided by the actual volume that was injected and finally multiplied by 100. The average of these residual values gave the linearity in percentage terms of individual data points for each specific compound. Analytical precision of 10 consecutive analyses of the same standard was better than 5% for all compounds except for methyl bromide (7%) and methyl chloroform (7%). We note that these relatively high values for methyl bromide and methyl chloroform were due to outliers. When omitting the extreme values (the lowest one and the highest one), then the precision is also better than 5% for these two compounds. We estimate the overall uncertainty of the measurements to be better than 15% for all compounds.

3 Results and discussion

Table 1 presents the results obtained for the VOC measurements during the MINOS campaign. The third column gives the overall geometric mean and the corresponding standard deviation. The high standard deviations for the alkanes are due to the large variability of their mixing ratios between clean and polluted events (see Sect. 3.2.1). In contrast, the CFCs which have longer lifetimes show less variability and hence lower standard deviations. For comparison with other studies, the average value observed on 7 August is included and indicated as “clean air” (Table 1). This day was the least influenced by recent pollution as indicated by the relatively low values of all measured hydrocarbons (e.g. alkanes and benzene). For the most reactive compounds, i.e. the hydrocarbons, a comparison with other measurements in the northern hemisphere is not straightforward as their concentrations are very variable depending on the proximity to, and intensity of their sources. Their variability in terms of sources and photochemical processing during MINOS are described in detail in Sect. 3.2.

For the long-lived compounds, the most extensive set of measurements is from the ALE/GAGE/AGAGE network (Prinn et al., 2000), which consists of 5 measurement stations in the northern hemisphere and 2 stations in the southern hemisphere. We compare our data to the measurements performed at Mace Head, the only ALE/GAGE/AGAGE station located in Europe. Five long-lived compounds (CFC-12, CFC-113, HCFC-142b, CHCl_3 and CCl_4) were measured both at Mace Head and at Finokalia during August 2001. The corresponding monthly means are reported in Table 1 for Mace Head and show a good agreement with the MINOS data. The highest difference is observed for CFC-12 between the MINOS selected clean air data and the Mace Head data. However, the campaign average agrees with the Mace Head August average within the uncertainty range.

3.1 Relationship of variability and lifetime of compounds

The lifetimes of the VOCs measured during MINOS vary between about 0.6 days (pentane) and 300 years (CFC-114). It has been shown previously that the variability of a compound and the measurement site are related to the sources and sinks of the compounds (Jobson et al., 1999; Jobson et al., 1998; Junge, 1974; Williams and al., 2000). Jobson et al. (1998) showed empirically that the variability and lifetimes of hydrocarbons can be linked through the equation $\text{Sln}X = A\tau^{-b}$. $\text{Sln}X$ represents the standard deviation of the logarithm of the mixing ratio of hydrocarbon X, τ is the chemical lifetime of the hydrocarbon and A and b are empirical fitting parameters. Jobson et al. (1999) extended this study by examining the variability/lifetime relationship for a larger number of VOCs (hydrocarbons and halocarbons). From these studies, it was proposed that the coefficient b, the dependence of variability on lifetime, reflects the importance of the source/sink budget

in the regional variability of the studied species and varies from $b=0$ being close to the sources to $b=0.5$ in remote locations. Since these first studies, the variability concept has been applied to a number of datasets and has proved useful in ascertaining the consistency and quality of measurements and for the estimation of OH concentrations (Williams et al., 2000; Williams et al., 2001, Karl et al., 2001).

The variability concept is based on the assumption that the chosen compounds have, more or less, the same source distribution. To a first approximation this is the case for the compounds reported in this paper, which are mostly of anthropogenic origin. Methyl chloride (chloromethane) and methyl bromide (bromomethane) are exceptions, as they have a globally important oceanic source (Brasseur et al., 1999). However, we note that no clear evidence for an ocean source or sink for either gas could be derived from the variability analysis of samples collected in the southern Pacific marine boundary layer (Colman et al., 1998).

The relationship between the variability and the lifetime of the VOCs measured during the MINOS campaign is shown in Fig. 1. The lifetimes of the reactive compounds were estimated by using the average measured OH value of $4.7 \times 10^6 \text{ molecules cm}^{-3}$ (Berresheim et al., 2003) and by assuming that OH is the only significant sink of these compounds. For the long-lived compounds, we used the global mean lifetimes as given by Jobson et al. (1999). A linear relationship of the form $\text{Sln}X = 0.85\tau^{-0.23}$ ($R=0.96$) was observed (Fig. 1). Note that the derived b coefficient is not sensitive to OH within the measurement uncertainty of OH ($\pm 30\%$). The fact that methyl bromide and chloride lie almost on the line may indicate that their anthropogenic emissions govern their variability in this region. However, this cannot be determined unequivocally from the variability plot.

The coefficient A, which relates to the range of air mass ages, is low (0.85), suggesting a narrow distribution of air mass ages. This is consistent with the range of the air mass ages estimated with toluene/benzene ratios (15–35 h, Salisbury et al., 2003) and ethylbenzene/acetophenone ratios (15–59 h, Xu et al., 2003). The coefficient b of this study has an absolute value of 0.23 and is significantly lower than the value of ~ 0.5 found during PEM-West B and at Amsterdam Island in the Indian Ocean (Jobson et al., 1999; Williams et al., 2001) but higher than the value of 0.18 found for the Harvard Forest site (Jobson et al., 1999). This suggests that Finokalia represents intermediate conditions between remote sites (where the variability is strongly dependent on the lifetime) and sites in the vicinity of sources (where sources dictate the variability, not the chemistry). In addition, the relatively quite high scatter observed on the plot suggests a wide variety of sources which are discussed further below. This is consistent with the back-trajectory analysis, which showed that several regions in Europe, and particularly in eastern Europe, influenced the air composition of Crete during the MINOS experiment (Salisbury et al., 2003).

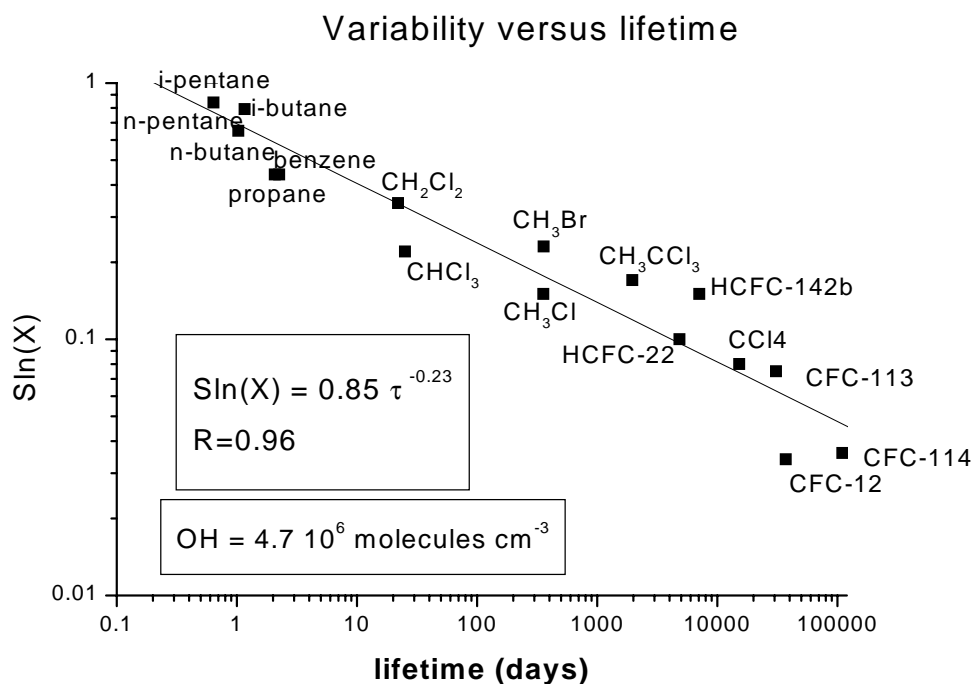


Fig. 1. Variability (standard deviation of $\ln[\text{mixing ratio}]$) against lifetime for the VOCs measured during the MINOS campaign.

In order to characterize the anthropogenic sources influencing the observations during MINOS better, the next sections will be dedicated to more detailed analyses of some specific compounds.

3.2 Fossil fuel and biomass combustion

3.2.1 Temporal variations of propane

Figure 2 presents the propane volume mixing ratios observed during the MINOS campaign (the propane values calculated with the global model MATCH are also presented on this graph and will be discussed in the next section). The classification of time periods is based on the variations of CO, black carbon (BC) and other compounds, and on back-trajectory calculations. More details about the definition of these periods can be found in Salisbury et al. (2003) and only a brief summary is given here. During period 1 (no VOC measurements during this period), air originating in Eastern Europe was observed. During period 2, the air originated above the boundary layer in Western Europe and was advected approximately east-south-east before descending and turning south-west over the Black Sea and north-western Turkey. Period 3 corresponds to a highly polluted event, mainly due to biomass burning emissions, originating from eastern Europe. Finally, period 4 corresponds to mixed trajectory origins, associated with some regional influence.

Although some scatter is associated with the data, the observed propane variations reflect clearly the air mass changes between periods. In period 2, the lowest values of propane

(~ 120 pptv) were observed. This is consistent with photochemically aged air descending from the free troposphere without recent contact with pollution sources. During period 3, significantly enhanced propane mixing ratios (up to 500 pptv) were observed, consistent with the corresponding high levels of CO and BC which have been attributed to the influence of biomass burning by reference to acetonitrile. We note that methyl chloride and benzene, two compounds emitted by biomass burning, presented large enhancements (up to 40% for methyl chloride and to a factor of 6 for benzene) during this period. The transition between period 3 and 4 was characterized by average levels of propane (~ 200 pptv) which reached high concentrations (up to 750 pptv) during period 4, consistent with some regional influence which was also evident from other compounds. Propane showed a large degree of scattering during period 4, reflecting the mixed origin of the air during that time.

3.2.2 Comparison with the global model MATCH

Figure 2 also shows the propane mixing ratios calculated by the MATCH-MPIC chemistry-transport model (Model of Atmospheric Transport and Chemistry – Max Planck Institute for Chemistry version). MATCH is a global atmospheric offline model, which can be driven by different meteorological datasets. Here the National Center for Environmental Prediction (NCEP) meteorological data at a horizontal resolution of about $2.8^\circ \times 2.8^\circ$ with 42 levels are used. MATCH is driven by meteorological parameters from the air mass trajectory re-analysis (Lawrence et al., 2003). The chemical

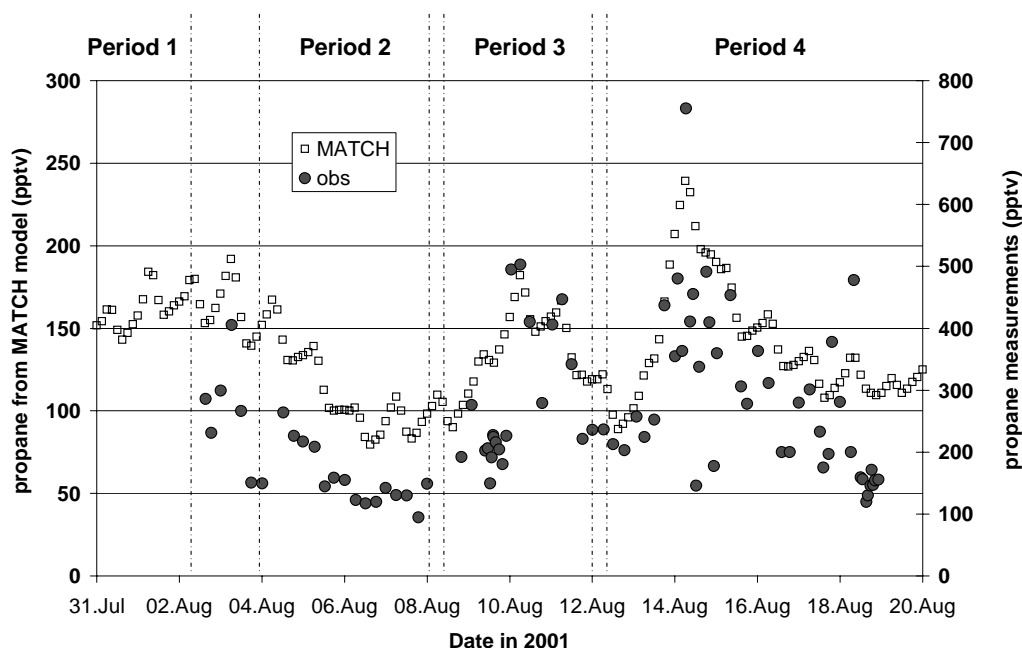


Fig. 2. Propane measurements during MINOS (closed circles, right-hand axis) and propane calculations from MATCH (open squares, left-hand axis). See text for the definition of periods.

scheme includes $\text{CH}_4\text{-CO-HO}_x\text{-NO}_x$ “background” chemistry, a simplified representation of isoprene chemistry, explicit ethane (C_2H_6) chemistry along with simpler representations of propane (C_3H_8), ethylene (C_2H_4), propylene (C_3H_6) and n-butane (C_4H_{10}) chemistry. The latter species is used as a surrogate for all higher alkanes. A complete description of the model can be found in Lawrence et al. (1999), von Kuhlmann (2001), von Kuhlmann et al. (2003a) and references therein and comparisons with the MINOS flight data are given in Lawrence et al. (2003). As propane is treated specifically, it is therefore the measured organic species most appropriate to directly compare with the model.

The model reproduces the main features observed during the campaign quite well, especially showing higher levels of propane during periods 3 and 4 and lower levels during period 2 and during the transition between period 3 and 4. However, the simulated mixing ratios of propane are lower than the observed data by a factor of 1.5 (background conditions) to 3 (polluted event observed in period 4).

The deviation between the modelled concentrations and our observations might have various causes, ranging from measurement errors, local influences in our measurements, erroneous modelling of eastern Mediterranean meteorology or inaccuracies in the emission data. We can rule out the first hypothesis, as the uncertainty in the measurements, estimated to be lower than 15% (Sect. 2.3), cannot account for the observed discrepancy between model and measurement. Local source influences did not cause the difference either; based on our interpretation of Fig. 1 (Sect. 3.1), the Finokalia site is hardly affected by local sources. Finally,

Table 2. Total NMVOC emissions in kt NMVOC in EDGAR database, except for biofuel and biomass burning emissions (source: v2.0, Olivier et al., 1996; v3.2, Olivier et Berdowski, 2001)

	V2.0 (1990)	V3.2 (1990)	V3.2 (1995)
Greece	244	291	312
Turkey	983	559	643
Ukraine	1980	7548	6736

errors in the model calculation are most likely not responsible for the discrepancy between model and measurement results: the model captures well the main variations of propane observed during MINOS, which indicates that the meteorology and the chemistry used in the model are realistic. Also, the modelled CO is very well correlated ($r=0.73$) with the flight data (Lelieveld et al., 2002). Therefore, an underestimation of the propane emissions used in the model is the most likely reason for the discrepancy. We note that propane concentrations simulated with the MATCH model were already compared to propane observations at several locations and that a reasonable agreement was found for northern Europe while an underestimation by a factor of 2 was found over North America (von Kuhlmann et al., 2003b). The discrepancy between measurements and model is therefore regionally dependent, which is consistent with the hypothesis that the problem comes from the emissions database. We will now dedicate the next section to discuss the propane emissions used for the MINOS comparison.

3.2.3 Emissions of propane

The propane emissions used in the model calculations were compiled by von Kuhlmann et al. (2003a). Emissions from energy and industrial activities (except for biofuel emissions) were taken from the Emission Database for Global Atmospheric Research (EDGAR v2.0, Olivier et al., 1996; Olivier et al., 1999a). Biomass burning emissions (including biofuel) were included by von Kuhlmann et al. (2003a) based on emission factors presented by Andreae and Merlet (2001). These emission factors were used to calculate a propane/CO emission ratio (biomass burning) for each type of biomass which was used to scale the CO emission distribution of Galanter et al. (2000). Together with emissions from the energy/industry system (5.4 Tg), biomass burning emissions (1.7 Tg) account for 95% of the global budget of 7.4 Tg.

The discrepancy between model and observations is observed for the whole campaign, including the beginning of the campaign where biomass burning influence was very low (Salisbury et al., 2003). Therefore, we will focus on possible errors in the energy/industry emission estimates and we will discuss successively four points: i) total VOC estimates, ii) speciation of the VOCs, iii) seasonal variation and iv) importance of the year of the simulation.

- i) In EDGAR, the total NMVOC (non-methane volatile organic compound) emissions were calculated using an emission factor approach. This emission might be inaccurate due to poorly known emission factors or activity data. In Olivier et al. (1999b) the uncertainty of the NMVOC emissions is estimated to be “medium” (50%) for fossil fuel related emissions and “large” (>100%) for non fossil fuel emissions.
- ii) The calculation of propane emissions in EDGAR are based on the application of sector-fuel specific NMVOC profiles which assign the fraction of each of 25 different NMVOC compounds from the total NMVOC emission. An inaccurate speciation of the compounds could also lead to a wrong estimation of propane emission. However, a calculation performed on the ratios of butanes/propane and pentanes/propane observed during MINOS suggests that the speciation provided by EDGAR is consistent with our measurements (see Sect. 3.2.4). This would further suggest that butane and pentane emissions are also similarly underestimated in the EDGAR database.
- iii) A third possible source of inaccuracy in the emission figures is that the energy/industry emissions are provided as total annual emissions. This means that the total propane emission was distributed uniformly over the year so that temporal variations such as seasonal changes or particular emission events are not taken into account.

- iv) Finally, the MATCH model simulation was performed with EDGAR v2.0 emission values that are representative for the year 1990. A detailed propane emission inventory for more recent years is not available at the moment. The question arises if and to what extent emissions of propane within the study domain have increased over the period 1990–2001. An indication of a possible increase in propane emissions can be found when looking at the recently constructed EDGAR v3.2 emission inventory. This inventory presents for total NMVOCs (not speciated) an update of 1990 emissions and an estimate of 1995 emissions. For countries that have a large influence on the MINOS observations (Greece, Turkey and Ukraine) the NMVOC emissions according to EDGAR v2.0 and EDGAR 3.2 are presented in Table 2. The results for total NMVOCs show both a systematic error in the calculation of Ukrainian emissions in version 2.0 and that emissions for more recent years (1995) are a factor 3.5 higher than the Ukrainian emissions used in the model calculations. For Greece and Turkey some small difference can be found. If this difference in total NMVOC emissions is also found in the propane emissions, then the discrepancy between model and observation can probably be explained by an underestimation of propane emissions in the Ukraine and maybe in other countries from the former Soviet Union. This could not be verified, since EDGAR v3.2 propane emissions are not yet available. The discrepancy between v3.2 and v2.0 for the year 1990 can largely be explained by the fact that in v2.0 Ukrainian emissions were calculated based on statistics from the total former Soviet Union, which were scaled to sub-regions based on e.g. population distributions. However, statistics have been revised in the past few years, new sources have been added and emission factors have been revised for some sources. During the construction of v3.2, country specific energy and industry statistics were available for the Ukraine (J. G. J. Olivier, personal communication).

3.2.4 Photochemical processing of the alkanes observed during MINOS

Assuming Lagrangian transport and that the only process affecting the ratio butane/propane (pentane/propane) since emission is the reaction with OH, one can easily derive the evolution of these ratios with time, as follows:

$$[A]_F/[B]_F = [A]_S/[B]_S \times \exp((k_B - k_A) \times [\text{OH}] \times \Delta t) \quad (1)$$

with

[A] mixing ratio of the compound A, the subscripts F and S stand for Finokalia and emission source region, respectively.

k_A : reaction rate of the compound A with OH

[OH]: OH concentration

Δt : transport time

Table 3. Butane^a/propane^b and pentane^c/propane^b ratios at the emission point (EDGAR database v2.0, Olivier et al., 1996), after chemical processing (see text) and observed at Finokalia.

		source	After 24 h	After 48 h	Finokalia
Greece	butane/propane	4.4	3.0	2.0	0.7±0.2
	pentane/propane	5.5	2.1	0.8	0.4±0.2
Turkey	butane/propane	1.7	1.2	0.8	0.7±0.2
	pentane/propane	2.0	0.8	0.3	0.4±0.2
Ukraine	butane/propane	1.6	1.0	0.7	0.7±0.2
	pentane/propane	1.6	0.6	0.2	0.4±0.2

^a $k_{\text{OH}}=2.26 \cdot 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, Donahue et al. (1998)

^b $k_{\text{OH}}=1.09 \cdot 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, Donahue et al. (1998)

^c $k_{\text{OH}}=3.92 \cdot 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$, Donahue et al. (1998), Baulch et al. (1986)

Ratios of butanes/propane and pentanes/propane were considered for this calculation. For comparison with the categories defined in the EDGAR database, iso and n-butane were aggregated in butanes and iso- and n-pentane in pentanes and the average reaction rate was considered (Baulch, 1986; Donahue et al., 1998). Emissions of the alkanes were extracted from the EDGAR database v2.0 (Olivier et al., 1996) for Greece, Turkey and Ukraine, three countries which likely had the most influence on the observations performed during MINOS. The values reported for Finokalia represent the average of the ratio (and its standard deviation) for the first part of the campaign (until end of period 2). Only the first part of the campaign has been considered in order to exclude periods which were affected by different emissions (nature of the emission and/or geographical origin of the emission). The concentration of OH was taken as the mean value measured during periods 1 and 2 ($4.0 \times 10^6 \text{ molecules cm}^{-3}$, Berresheim et al., 2003). Finally, transport times of 24 h and 48 h were considered to cover the range estimated by using toluene/benzene (Salisbury et al., 2003) and ethylbenzene/acetophenone ratios (Xu et al., 2003) for the corresponding periods.

Butane/propane and pentane/propane ratios at the emission point, after chemical processing (24 h and 48 h) and observed at Finokalia are shown in Table 3. Keeping in mind the limitations of this rough calculation, one can note that emissions from Turkey and Ukraine probably influenced the observations made during the first part of MINOS most strongly. Indeed, both observed ratios (butane/propane and pentane/propane) are consistent with emissions from these countries and transport times ranging from 24 h to 48 h.

Comparing our observations with the few other measurements of hydrocarbons performed in non-urban areas of Greece shows that the mixing ratios were in the same absolute concentration range for the C3–C5 hydrocarbons (Bonsang et al., 1999, Moschonas, 2000). However, we note that the VOC ratios were different in all three campaigns, reflecting different emission sources (the other studies were per-

formed over continental Greece, and were certainly more affected by Greek emissions) and different extents of photochemical processing.

3.3 Anthropogenic solvent emission

3.3.1 Temporal variation of methyl chloroform

Methyl chloroform (1.1.1. trichloroethane) has been widely used over the last 40 years as a solvent. However, its capacity to release chlorine in the stratosphere (leading to ozone destruction) has led to the phasing out of its production. This phasing out, initially planned for 2005 (Montreal protocol and London amendment), has been brought forward to 1996 for developed countries (Copenhagen amendment; McCulloch and Midgley, 2001). Therefore, methyl chloroform (MCF) emissions are now considered to be almost zero for mid-latitude countries in the northern hemisphere. The measured methyl chloroform mixing ratios during MINOS are shown in Fig. 3 and show large peaks on 5, 9 and 14 August. Since biomass burning is a small source of MCF (Rudolph et al., 2000), the observed simultaneous enhancements of CFC-113 indicate that other anthropogenic sources caused the MCF peaks observed during MINOS (Fig. 3). The good correlation ($R^2=0.5$ for the whole campaign) observed between MCF and CFC-113 ($\text{CCl}_2\text{FCClF}_2$), whose applications are similar to those of methyl chloroform, clearly indicate that significant emissions of these compounds still occur in Europe.

These peaks cause the relatively high variability of MCF, which pulls its value above the linear fit line in Fig. 1. If emissions had ceased, MCF should be below the line defined by species that still have source-generated variability, as was observed in data collected in the remote southern Indian Ocean (Warneke and de Gouw, 2001).

Finally, one should note that the increase of methyl chloroform and CFC-113 observed during period 3 clearly indicates that the large pollution event attributed to biomass burning emissions (Salisbury et al., 2003) also included some other

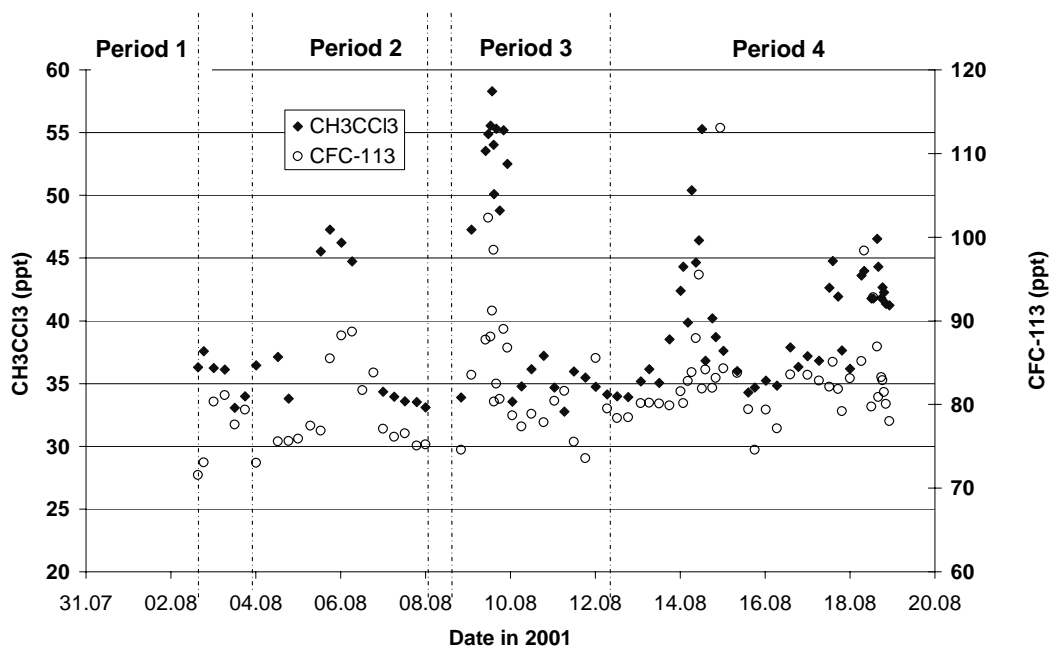


Fig. 3. Methyl chloroform and CFC-113 mixing ratios during MINOS.

anthropogenic emissions, at least on 9 August. In addition, a peak in propane was observed on 14 August which was not the case on 9 August. This likely indicates that the two events did not have the same source origin, which suggests that the significant methyl chloroform enhancements on 9 and 14 August may have originated in different countries.

3.3.2 Emissions strength of methyl chloroform

To evaluate the origin of the methyl chloroform emissions, we used an atmospheric-tracer transport model (Tracer Model, version 5: TM5) which allows studying the European region with a horizontal resolution of 1° by 1° . Vertically, 25 layers are used, which are mostly located in the boundary layer and in the free troposphere. The reference simulation of the model considers a total emission of 1 Tg/year methyl chloroform for Europe. Sensitivity simulations were performed by considering an additional methyl chloroform source (equivalent to 10 Tg/year) for the different European countries whose emissions may have influenced observations in Crete (Krol et al., 2003). The only exception to this procedure was for continental Greece, where the enhancement was only of 1 Gg/year (otherwise its proximity to Finokalia would lead to an unrealistically large impact). The results shown in Fig. 4 demonstrate that the main source of MCF observed at Finokalia on the 5 August could have been from Turkey, with possible contributions from Bulgaria and Greece, while the 9 August event corresponds to possible impacts from both Turkey and Ukraine (this last country signed the Copenhagen amendment in April 2002). The increase in methyl chloroform observed on 14 August could be attributed to Ukraine,

but also to Bulgaria or Romania. It should be noted that contributions from the Caucasus region (not included in this simulation), where high levels of MCF have been reported previously (Folberth et al., 2000, 2003), cannot be excluded.

3.3.3 Implication for OH calculation

Methyl chloroform has received much attention recently because its known anthropogenic emissions could help to indirectly determine the global mean OH concentration and trend (Spivakovsky et al., 2000; Prinn et al., 2001; Krol and Lelieveld, 2003 and references therein). Recently, Prinn et al. (2001) derived a strong negative trend of OH over the last decade, based on the assumption that emissions of MCF have ceased. However, during MINOS and other experiments performed in Europe (Krol et al., 2003), it has been shown that significant methyl chloroform emissions still take place in some parts of Europe. Therefore, the uncertainty associated with the current MCF emissions makes any conclusion on OH distribution and trend difficult. In particular, the previously derived trend is probably exaggerated and tropospheric OH may have been relatively constant over the last decade (Krol et al., 2003).

It is nevertheless important to note that these emissions (MCF and CFC-113) are still relatively small and that they will not have a significant adverse effect on the predicted recovery of stratospheric ozone.

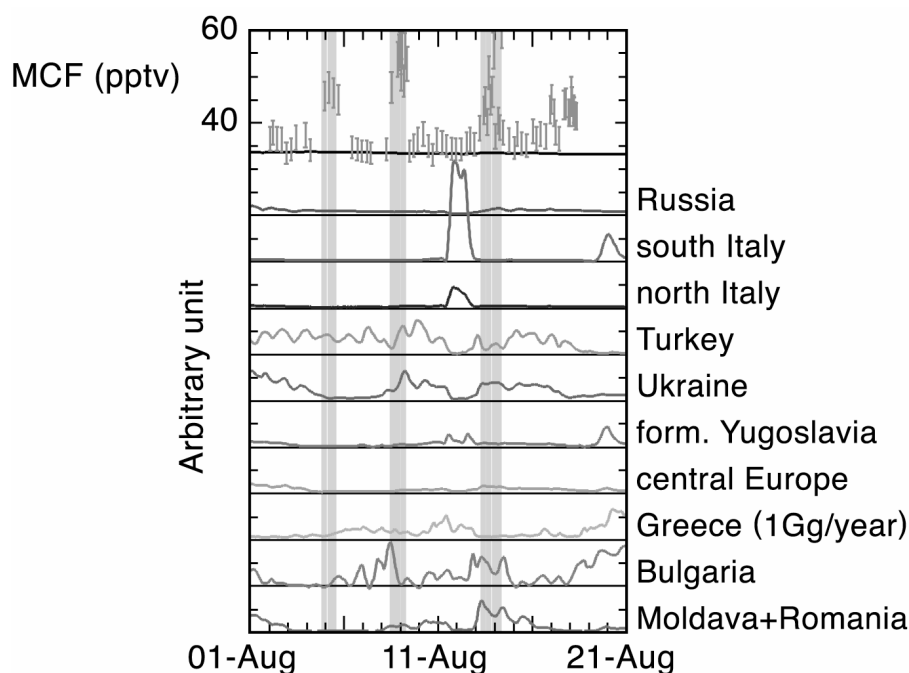


Fig. 4. Top panel: Methyl chloroform observations during MINOS and reference simulation of the model (black line going through the lowermost observations). Lower panels: sensitivity simulations for methyl chloroform emissions for different European regions (see text for details).

3.4 Anthropogenic agricultural emissions (methyl bromide)

The methyl bromide (CH_3Br) measurements are shown in Fig. 5 together with the previously discussed propane results. Along with its natural source from the ocean, methyl bromide is emitted by biomass burning, produced industrially and used for agricultural purposes (Schauffler et al., 1999 and references therein). Fumigation of soils before planting is the largest single use of CH_3Br worldwide (over 80%, (Lee-Taylor et al., 1998)).

The first point to note from Fig. 5 is that the methyl bromide mixing ratios observed on 7 August (defined as a “clean” day) were slightly higher than the overall mean (see Table 1). This is due to the definition of the “clean” period, which was chosen as the day the least influenced by recent pollution based on the low values of fossil fuel derived hydrocarbons measured that day. Due to the longer lifetime of methyl bromide, some older polluted air masses could still contain elevated concentrations of CH_3Br . Alternatively, the fact that CH_3Br has specific anthropogenic sources resulting from agricultural activity could also explain this feature.

Second, although biomass burning is estimated to contribute about 15% to the global budget of CH_3Br (Lee-Taylor et al., 1998), no significant enhancement of this compound was observed during the polluted event of period 3 where biomass burning influences predominated (Sect. 3.2). We note that considerable variability in the emission ratios of

CH_3Br emitted by biomass burning processes has been reported in the literature, depending strongly on the bromine content of the fuel and on the proportion of flaming to smoldering combustion (Andreae and Merlet, 2001). However, even if we consider the upper limit of published CH_3Br emission factors versus CO (Blake et al., 1996 and references therein), the 100 ppbv CO enhancement observed during this episode (Salisbury et al., 2003) would lead to a CH_3Br enhancement of about 1 pptv. This value, which represents the maximum expected impact, is only slightly higher than our analytical precision, which would explain why there is no significant enhancement of methyl bromide during the biomass burning episode.

Finally, although most of the measurements lie between 10 and 15 pptv, Fig. 5 shows that one noticeable peak of CH_3Br was observed on 12 August, with two values higher than 25 pptv, which then progressively decreased to 13 pptv (16 hours after the maximum). This peak is significant with respect to instrumental uncertainty and indicates that this compound was being emitted at this time. As no other measured VOC showed a concomitant increase, this peak of methyl bromide likely reflects the influence of a specific source of this compound. According to Fig. 4, which shows the sensitivity of the measurements to the emission regions, a large contribution from south/north Italy was observed on 12 August. As Italy is the largest consumer of CH_3Br as soil fumigant in Europe and the third most important worldwide (ICF, 1997 and references therein), it is plausible that this

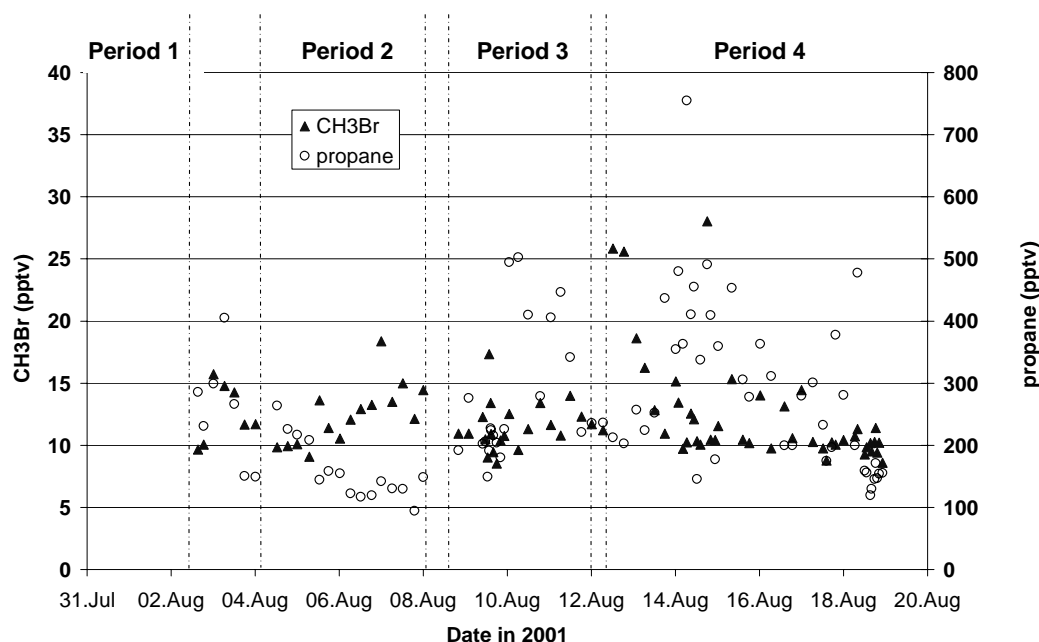


Fig. 5. Methyl bromide and propane mixing ratios during MINOS.

specific source of methyl bromide was responsible for the higher levels observed on the 12 August. While we cannot completely rule out the contribution of other specific sources of methyl bromide (such as salt marshes), we conclude that fumigant emissions from Italy are the most probable.

4 Summary and conclusions

The MINOS campaign provided a good opportunity to investigate the anthropogenic origin of several VOCs from European sources, in particular from southern and eastern Europe. By studying the variations observed during the campaign, we have derived information about the anthropogenic emissions of propane, methyl chloroform and methyl bromide. Propane, mainly emitted by fossil fuel and industrial processes, was compared with the simulated propane from the global model MATCH-MPIC. While the model reproduces the general features of propane variations quite well, it underestimates significantly its mixing ratios, which is attributed to an underestimation of the emissions (provided by the global database EDGAR). Initial investigations suggest that the total amount of VOCs accounted for in the database in some countries is too low, but further research is needed to address this issue. Emission inventories urgently need updating if models are to be usefully compared with measurements.

Special attention has also been dedicated to the interpretation of the methyl chloroform measurements, as some unexpected deviations from the background were observed during MINOS. This observation, complementing other measure-

ments in Europe, shows that methyl chloroform emissions still occur in Europe. These emissions need to be precisely quantified if the method of OH estimation (distribution and trend) based on methyl chloroform is to be further applied.

The methyl bromide measurements indicated that no significant regional biomass burning sources were detected. Variations of CH₃Br were not associated with fossil fuel usage but with a specific source region (Italy) and most probably associated with agricultural fumigant usage.

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